Table 1: Comparison of diagnostic methods for SMA

N° tube	SSCP ⁽¹⁾	Our méthod (radioactive) (3)							
	j	Probe 1 (exon 7)			Probe 2 (exon 8)				
	Exons (2)	PSL/mm ²	R (%) (4)	Exon 7	PSL/mm ²	R (%) (4)	Exon 8		
Control	ndel 7/ndel 8	42	0	ndel	22	0	ndel		
(1)				[
Control	ndel 7/ndel 8	41	0.	ndel	23	0	ndel		
(2)				<u> </u>					
Control	ndel 7/ndel 8	43	0	ndel	22	0	ndel		
(3)						<u> </u>			
Control	ndel 7/ndel 8	41	0	ndel	21	0	ndel		
(4)	<u> </u>								
Control .	ndel 7/ndel 8	42	0	ndel	23	0	ndel		
(5)			ļ .		ļ				
SMA	del 7/del 8	24	43	del	15	32	del		
(6)				<u> </u>					
SMA	del 7/del 8	14	67	del	08	64	del		
(7)	4-17/4-10	10	77.	-2-1	0.7				
SMA	del 7/del 8	10	76	del	07	68	del		
(8) \$MA	del 7/del 8	26	38	del	06	73	del		
3MA (9)	dei //dei o		30	Gei		13	dei		
SMA	del 7/del 8	09	79	del	13	41	del		
(10)	derivation	Ü	13	u c.		•	40.		
SMA	del 7/del 8	27	36	del	12	45	del		
(11)					,_	, ,			
SMA	del 7/del 8	15	64	. del.	15	32	del		
(12)									
SMA	del 7/del 8	13	69	del	14	36	del		
(13)						,			
SMA	del 7/del 8	25	40	del	11	50	del		
(14)									
SMA	del 7/del 8	20	52	del	.09	59	del		
(15)		, , ,							
SMA	del 7/del 8	19	55	del	13	41	del		
(16)			~ .						
SMA	del 7/del 8	12	71	del	14	36	del		
(17)	2017/210			4 - 1			٠		
SMA	del 7/del 8	14	67	del	12	45	del		
(18)	rand conformatio	<u> </u>					[

⁽¹⁾ single strand conformation polymorphism(2) del: deleted; ndel: non deleted

The mean value of the control group is used for the calculation of R

⁽³⁾ The quantification of results obtained is performed by means of Bio-Imager and expressed as PSL/mm²

⁽⁴⁾ R: difference = 1 - [(PSL/mm²Control - PSL/mm²SMA)/(PSL/mm²Control)]

Table 2: Comparison of diagnostic methods for SMA

N° tube	SSCP (1)	Our method (ELISA)								
			Deals 1 /cm	on 71	Probe 2 (exon 8)			Probe 3 (HUMEF1AB)		
		Optical	Probe 1 (exon 7) Optical Exon 7		Optical	rroue Z (ex	Exon 8	Optical	RT-PCR	
	Exons (2)	density	R(%) ⁽³⁾		density	R(%) ⁽³⁾		density		
Control	ndel 7/	0.26	0	ndel	0.28	0	ndel	0.55	positive	
(1)	ndel 8									
Control	ndel 7/	0.28	0	ndel	0.29	0	ndel	0.54	positive	
(2)	ndel 8			•						
Control	ndel 7/	0.27	0	ndel	0.26	0	ndel	0.52	positive	
(3)	ndel 8									
Control	ndel 7/	0.26	0	ndel	0.28	0	ndel	0.51	positive	
(4)	ndel 8									
Control	ndel 7/	0.27	0	ndel	0.27	0	ndel	0.49	positive	
(5)	ndel 8									
SMA	del 7/	0.16	41	ďei	0.13	54	del	0.51	positive	
(6)	del 8		4.4		0.10			0.10	•	
SMA	del 7/	0.16	41	del	0.16	· 43	del	0.48	positive	
(7)	dei 8					p		0.50	•.•	
SMA	del 7/	0.11	59	del	0.12	57	del	0.53	positive	
(8)	del 8 del 7/	015	44	del	0.14	50	del	0.49		
SMA	del 8	0.15	44	aeı	0.14	30	Gei	0.43	positive	
(9) SMA	del 7/	0.19	30	del	0.16	43	del	0.5	positive	
(10)	del 8	0.15 .	30	ue,	0.10	42	Gei	0.5	positive	
SMA	del 7/	0.13	52	del	0.12	57	del	0.49	positive	
(11)	del 8	0		00,	V., Z.	*	J G.	00	positivo	
SMA	del 7/	0.14	48	del	0.13	54	del	· 0.48	positive	
(12)	del 8				1	- 1				
SMA	del 7/	0.17	37	del	0.14	50	del	0.55	positive	
(13)	del 8									
SMA	del 7/	0.12	55	del	0.15	46	del	0.52	positive	
(14)	del 8			<u> </u>						
SMA	del 7/	0.11	59	del	0.13	54	del	0.5	positive	
(15)	del 8		l							
SMA	del 7/	0.13	52	dei	0.15	46	del	0.51	positive	
(16)	del 8									
SMA	del 7/	0.15	44	del ·	0.12	57	del	0.49	positive	
(17)	del 8									
SMA	del 7/	0.16.	41	del	0.16	43	del	0.5	positive	
(18)	del 8					1				

⁽¹⁾ single strand conformation polymorphism

The mean value of the control group is used for the calculation of R

⁽²⁾ del: deleted; ndel: non deleted (3) R: difference = $1 - [(OD_{450}Control - OD_{450}SMA/OD_{450}Control)]$